Nanoporous/mesoporous palladium catalyst

#### **PRIORITY**

[0001] This Application claims the benefit under 35 U.S.C. § 371 of International Application No.: PCT/GB2005/000002 (published PCT application No. WO 2005/067082), filed January 5, 2005, which claims priority to G.B. Patent Application No.: 0400166.5 filed January 6, 2004; the entire contents of each of the above-referenced applications are incorporated herein by reference.

### FIELD OF THE INVENTION

<u>100021</u> The present invention relates to a catalytic system comprising a catalyst comprising nanoporous or mesoporous palladium and an ion-exchange electrolyte. The invention also provides a multi-component catalyst which comprises nanoporous or mesoporous palladium and one or more additional components. Also provided are processes for manufacturing the catalytic system and catalyst, and a process for oxidising or reducing organic and/or inorganic molecules using said catalyst or catalytic system. The invention further relates to an electrode comprising the catalyst or the catalytic system, and also a fuel cell or a sensor comprising the catalyst, the catalytic system or the electrode.

### **BACKGROUND OF THE INVENTION**

[0003] There has been much interest in the development of fuel cells and sensors in the past decades. The development of such fuel cells and sensors has been highly dependent upon the use of catalysts for the oxidation of H<sub>2</sub>, small organic molecules and hydrocarbons, as well as for the reduction of O<sub>2</sub>. Catalysts for the oxidation of molecules such as H<sub>2</sub>, small organic molecules and hydrocarbons, etc are typically noble metals in dispersed form. For the reduction of oxygen, the most active catalyst is considered to be one composed of PtCr spherical or cubo-octahedral particles supported on carbon, (T.R. Ralph and M.P. Hogarth, *Platinum Metals Review*, 2002, 46, (1), 3-14) whereas for pure hydrogen, the most active catalyst is Pt spherical or cubo-octahedral particles also supported on carbon. For the oxidation of reformate (hydrogen containing some carbon monoxide), the preferred catalyst is one composed of PtRu or PtMo spherical or cubo-octahedral particles supported on carbon particles (T.R. Ralph and M.P. Hogarth, *Platinum Metals Review*, 2002, 46, (3), 117-135) whereas for the oxidation of methanol, and similar small organic molecules, the preferred catalyst is PtRu alloy in the form of

small particles either unsupported or supported upon carbon particles (T.R. Ralph and M.P. Hogarth, *Platinum Metals Review*, 2002, **46**, (4), 146-164).

[0004] Within the current art, typical electrodes consist of a gas diffusion substrate with platinum-based (Pt) electrocatalyst layers located between the ion-exchange membrane electrolyte and the substrate. The electrocatalyst can be deposited either on the gas diffusion substrate or on the ion-exchange membrane electrolyte using techniques such as screen printing, flexographic printing, gravure printing, spraying or rolling and calendering. Electrocatalyst layers are typically from 5 to 20 μm thick and the layers are normally bonded together by hot pressing catalysed substrates to the membrane or, in the case of catalysed membranes, by compressing the gas diffusion substrate to the membrane (T.R. Ralph and M.P. Hogarth, *Platinum Metals Review*, 2002, 46, (1), 3-14).

[0005] Virtually all catalysts for low temperature polymer electrolyte fuel cells (PEMFC) or electrochemical sensors utilise platinum as the major catalyst component, optionally alloyed with other species. Palladium has been considered as an alloying component but the performance of a platinum catalyst containing palladium based on current production methods is exceedingly poor, requiring 300mV extra overpotential to function at the same rate as a PtRu catalyst. As a result, such palladium containing catalysts would not normally be considered suitable for use as an electrocatalyst in a fuel cell or sensor.

<u>100061</u> In these previous cases, the catalysts exist in the form of fully dense nanoparticles – i.e. particles with a spherical or cubo-octahedral morphology with diameters ranging from about 1nm to 10 nm, possibly deposited on an electrically conducting but catalytically inactive support (e.g. carbon). The small dimensions of these particles are necessary to have a high surface area to volume ratio. An alternate structure to this involves catalyst materials which are not fully dense, but rather which have a mesoporous or nanoporous structure. Such materials may have a range of morphologies – e.g. spherical, cylindrical, film-like – but rather than being fully dense, they are porous and thus can still have a high surface area to volume ratio, even though the primary particle size is much larger than the fully dense particles currently used as catalysts. Examples of methods for making nanoporous or mesoporous materials include using templates such as a surfactant or mesoporous silicon. Other examples involve the use of a sol-gel synthesis route, the use of microemulsions or the use of metallo-organic chemical vapour deposition (MOCVD).

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[0007] As is described in this invention, for certain specific morphologies of palladium and

palladium containing catalysts, and in contact with specific types of electrolytes, the performance

of these catalysts may be significantly higher than might be expected by comparison to catalytic

systems containing palladium and/or palladium containing catalysts produced using standard

approaches.

**SUMMARY OF THE INVENTION** 

[0008] In a first aspect, the present invention provides a catalytic system comprising a catalyst

comprising nanoporous or mesoporous palladium and an ion-exchange electrolyte.

[0009] In certain embodiments of the first aspect, the catalyst further comprises one or more of

platinum, gold, ruthenium, rhodium, osmium, iridium, silver, nickel, copper, cobalt, iron,

chromium, lead, vanadium or tungsten. In certain other embodiments, the catalyst is Pd, PdPt,

PdAu, PdPtRu, PdPtIr, PdPtAu, PdPtRuIr, PdPtRuOs or PdPtRuIrOs.

[0010] In certain other embodiments of the first aspect, the ion-exchange electrolyte is a cation

exchange electrolyte.

[0011] In a second aspect, the present invention provides a multi-component catalyst comprising

a nanoporous or mesoporous palladium and one or more of platinum, gold, ruthenium, rhodium,

osmium, iridium, silver, nickel, copper, cobalt, iron, chromium, lead, vanadium, tungsten,

carbon, nitrogen, oxygen, sulphur, selenium, tellurium or phosphorous.

[0012] In a third aspect, the present invention provides a process for the production of a catalyst

of the second aspect of the invention, wherein the process comprises solidifying one or more

metal precursors in the presence of a templating agent and then removing the templating agent.

[0013] In a fourth aspect, the present invention provides a process for the oxidation or reduction

of inorganic and/or organic molecules comprising contacting the molecules with a catalytic

system of the first aspect of the invention, or a multi-component catalyst of the second aspect of

the invention.

[0014] In certain embodiments of the fourth aspect, the organic molecule has from one to twelve

carbon atoms. In certain other embodiments, the organic molecule is methanol.

[0015] In certain embodiments of the fourth aspect, in the catalytic system, the catalyst is Pd,

PdPt, PdAu, PdPtRu, PdPtIr, PdPtAu, PdPtRuIr, PdPtRuOs or PdPtRuIrOs. In certain

embodiments, in the catalytic system, the ion-exchange electrolyte is a cation exchange

electrolyte.

[0016] In a fifth aspect, the present invention provides a process for the manufacture of a

catalytic system of the first aspect of the invention, which process comprises contacting the

catalyst with the ion-exchange electrolyte.

[0017] In certain embodiments of the fifth aspect, in the catalytic system, the catalyst is Pd,

PdPt, PdAu, PdPtRu, PdPtIr, PdPtAu, PdPtRuIr, PdPtRuOs or PdPtRuIrOs. In certain other

embodiments, in the catalytic system, the ion-exchange electrolyte is a cation exchange

electrolyte.

[0018] In a sixth aspect, the present invention provides an electrode comprising a catalytic

system of the first aspect of the invention, or a multi-component catalyst of the second aspect of

the invention.

[0019] In certain embodiments of the sixth aspect, in the catalytic system, the catalyst is Pd,

PdPt, PdAu, PdPtRu, PdPtIr, PdPtAu, PdPtRuIr, PdPtRuOs or PdPtRuIrOs. In certain other

embodiments, in the catalytic system, the ion-exchange electrolyte is a cation exchange

electrolyte.

[0020] In a seventh aspect, the present invention provides a fuel cell comprising an electrode of

the sixth aspect of the invention.

[0021] In certain embodiments of the seventh aspect, in the catalytic system, the catalyst is Pd,

PdPt, PdAu, PdPtRu, PdPtIr, PdPtAu, PdPtRuIr, PdPtRuOs or PdPtRuIrOs. In certain other

embodiments, in the catalytic system, the ion-exchange electrolyte is a cation exchange

electrolyte.

[0022] In an eighth aspect, there is provided a sensor comprising an electrode of the sixth aspect

of the invention.

[0023] In certain embodiments of the eighth aspect, in the catalytic system, the catalyst is Pd,

PdPt, PdAu, PdPtRu, PdPtIr, PdPtAu, PdPtRuIr, PdPtRuOs or PdPtRuIrOs. In certain other

embodiments, in the catalytic system, the ion-exchange electrolyte is a cation exchange

electrolyte.

[0024] In a ninth aspect, there is provided a method for the detection of organic and/or inorganic

molecules in a sample comprising contacting said sample with a sensor of the eighth aspect of

the invention and detecting the current due to the oxidation or reduction of the molecules.

BRIEF DESCRIPTION OF THE DRAWING

[0025] Figure 1 shows SEM image and TEM image (inset) of Palladium mesoporous film

electrochemically deposited on a gold foil at 0.16 V vs Ag/AgCl for 4 hr with a charge of 20 C

 $cm^{-2}$ .

[0026] Figure 2 shows high resolution scanning electron micrograph images of PtPdRuOs alloy

chemically prepared from a liquid crystal phase, (A). Transmission electron micrograph (TEM)

images of PtPdRuOs nanoparticles, (B). TEM images of a single PtPdRuOs particle, (C).

Electron diffraction patterns of aggregated PtPdRuOs nanoparticles (D)

[0027] Figure 3 shows cyclic voltammograms for a mesoporous (a) and polycrystalline (b)

Palladium disc microelectrode in 0.1 mol dm<sup>-3</sup> HClO<sub>4</sub> + 1 mol dm<sup>-3</sup> methanol (dash line) and on

a Nafion<sup>TM</sup> membrane equilibrated with 1 mol dm<sup>-3</sup> methanol in water (solid line) at 20 °C

 $(dE/dt: 50 \text{ mV s}^{-1}):$ 

[0028] Figure 4 shows steady-state voltammograms for microelectrodes composed of

mesoporous palladium, mesoporous PdPtRu and mesoporous PdPtRuOs catalysts on a Nafion<sup>TM</sup>

membrane equilibrated with 1 mol dm<sup>-3</sup> CH<sub>3</sub>OH in water (a); and for mesoporous palladium on a

Nafion<sup>TM</sup> membrane equilibrated with 0.15 mol dm<sup>-3</sup> HCOOH in water and 0.20 mol dm<sup>-3</sup>

HCHO in water, (b).

[0029] Figure 5 shows steady-state voltammograms measured at 5 mV s<sup>-1</sup> for a microelectrode

containing mesoporous PtPdRuOs catalyst on a Nafion membrane equilibrated with 1 mol dm<sup>-3</sup>

methanol in water at various temperatures. Inset: Potential dependence of mass activity of the

quaternary alloy at 20 and 60 °C. Data taken from the positive-going scans.

[0030] Figure 6 shows a plot of limiting current versus methanol concentration (in water) for a

mesoporous palladium catalyst/Nafion<sup>TM</sup> membrane catalyst system.

DETAILED DESCRIPTION OF CERTAIN PREFERRED EMBODIMENTS OF THE INVENTION

[0031] The first aspect of In a first aspect, the invention herein described provides a catalytic

system comprising a catalyst comprising nanoporous or mesoporous palladium and an ion-

exchange electrolyte.

[0032] The In certain embodiments, the catalyst as described in this invention exists as a bulk

material that comprises palladium having a nanoporous or mesoporous morphology. It is these

pores that contribute to the large specific surface area of the catalyst. The pore size may range in

diameter from about 0.75nm to about 50nm, preferably about 0.75 to about 20nm, and more

specifically about 1 to about 10nm diameter. The distance separating the pores follows a similar

range.

[0033] The catalytic system of the first aspect provides improved performance compared with

previously reported systems containing palladium containing catalysts.

[0034] It is hypothesised that the improvement in activity of the catalytic system of the present

invention over other palladium containing catalytic systems is related to an alteration in the

formation of adsorbed oxygen-containing species on the catalyst. This modification of adsorption

properties may be the result of an interaction between the nanoporous or mesoporous structure of

the catalyst with the ion-exchange electrolyte.

[0035] In a preferred feature, the palladium catalyst comprises nanoporous or mesoporous

palladium with one or more additional components so as to form a multi-component catalyst.

Said multi-component catalyst can comprise one, two three, four or more components in addition

to palladium. Preferably the additional component can be a transition metal (e.g. a precious

metal), or a mixture of transition metals. Preferably the additional component or components are

alloyed with the palladium. Alternatively they can be present as a mixture with the palladium.

The additional component can be one or more selected from, platinum, gold, ruthenium,

rhodium, osmium, iridium, silver, nickel, copper, cobalt, iron, chromium, lead, vanadium or

tungsten. Alternatively, the component can be non-metallic, for example carbon, nitrogen,

oxygen, sulphur, selenium, tellurium, phosphorous. Preferably the additional component is

platinum and/or gold.

[0036] The In certain embodiments, the catalyst is preferably provided as a two component alloy

or mixture for example PdPt or PdAu, a three component alloy or mixture for example PdPtRu,

PdPtIr, PdPtAu, a four component alloy or mixture for example PdPtRuIr, PdPtRuOs and/or a

five component alloy or mixture for example PdPtRuIrOs. The catalyst may comprise one or

more of the two, three, four or five component alloys or mixtures.

[0037] The amount of palladium present in relation to the other components can vary from about

10 % to about 100 % in atomic percentage. The amount of palladium present will depend upon

the purpose of the application of the catalytic system. When palladium is the main component,

its content will be very high (close to 100 At %). When a high amount of non-palladium

components are necessary for the activity of the catalytic system, then the amount of palladium

can be lower, for instance at a level of about 20 At % or less, for instance about 10 At%.

[0038] The catalyst comprising nanoporous or mesoporous palladium can be prepared by

solidification of a metal precursor in the presence of a templating agent.

[0039] The templating agent may be one of a number of different compounds or systems of

compounds. For example, the templating agent may involve:

a) a template which is a non-ionic surfactant and which forms a liquid crystalline phase

under suitable conditions. This liquid crystalline phase directs the deposition of the

catalyst, and the catalyst then represents a 'cast' of that liquid crystalline phase;

b) the use of a suitable surfactant system resulting in the formation of a micro-emulsion of

the reactants which then undergo reaction to produce an interconnected nanoporous or

mesoporous structure of the catalyst. For example the synthesis of the mesoporous

palladium by mixing two water in oil microemulsions with the same water-surfactant

ratio, one containing H<sub>2</sub>PdCl<sub>6</sub> and the other containing hydrazine hydroxide;

c) the use of a suitable surfactant and conditions so that the oxide of the catalyst material is

produced in a 'sol-gel' reaction. Amphiphilic poly(alkylene oxide) block copolymers

such as P-123 and Brij<sup>®</sup>56, can act as structure-directing agents in nonaqueous solutions

for organizing network-forming metal species. The resultant mesoporous metal oxides

can be further reduced to metal systems;

d) a "hard" template – that is a template which is itself a solid mesoporous or nanoporous

material. The voids of a preformed nano- or mesoporous solid, such as mesoporous silica

and nanoporous alumina, are impregnated with metal-containing precursor or precursors.

Subsequent solidification of these precursors and removal of the 'hard' template leads to

nano- or meso- porous catalysts. Such 'hard' templates may be produced using the

previously mentioned approaches (a-c) or may be purchased from a suitable supplier.

[0040] The In certain embodiments, the solidification process converts the metal precursor to a

chemical species which is a solid and which can self-support itself in the absence of the

templating agent. After this solidification process, the templating agent is removed by, for

example, washing the template out of the system.

[0041] The solidification process may involve a chemical transformation of a precursor by for

instance the oxidation or reduction of the metal precursor, or a change in the coordinating species

around the metal precursor with no change in formal oxidation state.

[0042] The reduction process can be carried out by any process known in the art by for example

(a) Electrochemically reducing – by poising the electrochemical potential of an electrode within

a suitable range so that the metal precursor is reduced to its corresponding metals on the surface

of the electrode; or (b) Chemically reducing – by adding a chemical reductant which reduces the

metal precursor to its corresponding metal. Suitable chemical reductants are for instance zinc,

iron, calcium or magnesium. Alternatively liquid reductants such as formaldehyde solutions;

formic acid solutions; or borohydride solutions; or gaseous reductants such as hydrogen,

formaldehyde, or boranes can be used.

[0043] Oxidation In certain embodiments, the oxidation process can be carried out by any

process known in the art for example (a) Electrochemically oxidising - by poising the

electrochemical potential of an electrode within a suitable range so that the metal precursor is

oxidised and forms an insoluble compound; or (b) Chemically oxidising – by adding a chemical

oxidant which oxidises the metal precursor to an insoluble compound. Suitable chemical

oxidants include solutions containing hydrogen peroxide or peroxide salts; perborates,

permanganates, chromates or chlorates; or gaseous oxidants such as ozone, chlorine or oxygen.

[0044] Another approach to cause solidification of the metal precursor involves the formation of

an insoluble chemical species. In one example, this can involve the addition of a compound

which changes the pH so that a metal oxide or hydroxide is precipitated. Compounds which can

be used to change the pH are acidic or basic materials, especially strong acids and bases for

instance sulphuric, nitric or hydrochloric acid or alkali metal hydroxides.

[0045] The metal precursor for the purposes of this invention should exhibit a high solubility in

the solution containing the templating agent. Such precursors can be the chloro complexes (i.e.

metal chlorides) or other halide complexes, or the amino (NH<sub>3</sub>), aqua (H<sub>2</sub>O), sulfito (SO<sub>3</sub>), or

cyano (CN) coordinated complexes, or complexes which contain a mixture of one or more of the

above ligands. These complexes are particularly preferred for use in aqueous systems.

Complexes soluble in non-aqueous solvents include complexes containing acetylacetonate or

hexafluoroacetylacetonate, 1,10-phenanthroline and derivatives thereof, 1,5-cyclooctadiene and

derivatives thereof, or ethylenediamine and derivatives thereof.

[0046] In the present invention, In certain embodiments, the catalyst is placed in contact with an

ion-exchange electrolyte. The ion-exchange electrolyte is preferably a cation exchange material

having fixed anionic sites. More preferably, the ion-exchange electrolyte is a perfluorinated

polymer electrolyte having a pendant acidic group such as a sulphonic acid group teathered to a

perfluorinated backbone. Examples of such perfluorinated polymer electrolytes include Nafion®

(Dupont), Aciplex<sup>®</sup> (Asahi Chemical Industry), Flemion<sup>®</sup> (Asahi Glass Company) or BAM<sup>®</sup>

(Ballard).

[0047] Other cation exchange electrolytes which can be used for the purposes of the present

invention are electrolytes with pendant acidic groups such as sulphonic or phosphonic acid

groups with a hydrocarbon polymer backbone. Examples of such electrolytes include sulfonated

or phosphonated forms of polyamide/imide (Kerimid, Torlon®); polychlorotrifluoroethylene,

polyetheretherketone (PEEK), polyethersulfone (PES), polyimide, or polyphenylenesulfide, or

derivatives thereof.

[0048] The ion-exchange electrolytes can also be provided as selected solid-state inorganic

proton-conductors, such as derivitised zirconium phosphonates, immobilised and derivatised

Keggin and Dawson acids (e.g. phosphomolybdic acid); or hot proton-conductors, such as

perovskite-type oxide proton-conducting ceramics.

[0049] The ion exchange electrolyte can be provided as a solid in a variety of forms (the bulk

electrolyte). For instance it can be provided as a sheet. Alternatively, the ion exchange

electrolyte can be provided in a liquid form (the liquid ionomer) which can then form the solid

electrolyte after a suitable treatment process. For instance the liquid ionomer may consist of the

ion exchange electrolyte material dissolved in some suitable solvent, or it may be the ion-

exchanged electrolyte. Alternatively, the liquid ionomer can be a chemical precursor to the said

ion-exchange electrolyte. Suitable processes for converting the liquid ionomer to the solid ion-

exchange electrolyte can be evaporation of the solvent, changing in temperature of the liquid,

exposure to radiation, initiation of cross-linking or polymerisation or a process known in the art.

[0050] An especially preferred ion-exchange electrolyte is a perfluorinated ion-exchange

polymer consisting of a polytetrafluoroethylene backbone and regular spaced long perfluorovinyl

ether pendant side chains terminated by a sulfonate ionic group (Nafion<sup>TM</sup>). One example of the

catalytic system provided herein is thus a catalyst comprising nanoporous or mesoporous

palladium in contact with Nafion<sup>TM</sup>.

[0051] The second aspect of the invention provides In a second aspect of the invention, there is

provided a process for the manufacture of a catalytic system in which a catalyst comprising

nanoporous or mesoporous palladium is placed in contact with an ion-exchange electrolyte, so

that the catalyst is in ionic contact with that electrolyte.

[0052] This ionic contact can result from one or more of

simple mechanical force; a)

the use of an adhesive to bind the catalyst to the electrolyte; b)

the use of a liquid ionomer form of the electrolyte to interpenetrate the porous c)

structure of the catalyst and to form a bulk phase of that electrolyte adjacent to

the catalyst after suitable treatment; or

the use of a liquid ionomer to interpenetrate the porous structure of the d)

catalyst and bind it to a sample of the bulk electrolyte after suitable treatment.

This liquid ionomer can be a soluble form of the bulk electrolyte or it may be

a liquid ionomer form of a different ion-exchange electrolyte.

[0053] The third aspect of the invention provides In a third aspect of the invention, there is

provided a multi-component catalyst comprising nanoporous or mesoporous palladium and one

or more of platinum, gold, ruthenium, rhodium, osmium, iridium, silver, nickel, copper, cobalt,

iron, chromium, lead, vanadium, tungsten, carbon, nitrogen, oxygen, sulphur, selenium, tellurium

or phosphorous. Preferably the multi-component catalyst comprises one or more transition

(precious) metals such as platinum, gold, ruthenium, rhodium, osmium, iridium, silver, nickel,

copper, cobalt, iron, chronium, lead, vanadium or tungsten, more preferably platinum and/or

gold.

[0054] The In certain embodiments, the multi-component catalyst is provided as a mixture of

palladium and one or more additional components, preferably as an alloy. The catalyst is

preferably provided as a two component alloy or mixture thereof for example PdPt or PdAu, a

three component alloy or a mixture thereof for example PdPtRu, PdPtLr, PdPtAu, a fair

component alloy or a mixture thereof, for example PdPtRuLr, PdPtRuOs and/or a five

component alloy or mixture thereof, for example PdPtRuLrOs. The multi-component catalyst

can comprise one or more of the two, three, four or five component alloys or mixtures.

[0055] The In certain embodiments, the catalyst comprises 10% to 100% atomic percentage,

preferably 20% to 100% atomic percentage.

[0056] The fourth aspect of the invention provides In a fourth aspect of the invention, there is

provided a process for the production of a multi-component catalytic system according to the

third aspect of the invention, said process comprising solidifying one or more metal precursors in

the presence of a templating agent and then removing said templating agent.

[0057] The templating agent may be one of a number of different compounds or systems of

compounds. For example, the templating agent may involve:

e) a template which is a non-ionic surfactant and which forms a liquid crystalline phase

under suitable conditions. This liquid crystalline phase directs the deposition of the

catalyst, and the catalyst then represents a 'cast' of that liquid crystalline phase;

f) the use of a suitable surfactant system resulting in the formation of a micro-emulsion of

the reactants which then undergo reaction to produce an interconnected nanoporous or

mesoporous structure of the catalyst. For example the synthesis of the mesoporous

palladium by mixing two water in oil microemulsions with the same water-surfactant

ratio, one containing H<sub>2</sub>PdCl<sub>6</sub> and the other containing hydrazine hydroxide;

g) the use of a suitable surfactant and conditions so that the oxide of the catalyst material is

produced in a 'sol-gel' reaction. Amphiphilic poly(alkylene oxide) block copolymers

such as P-123 and Brij<sup>®</sup>56, can act as structure-directing agents in nonaqueous solutions

for organizing network-forming metal species. The resultant mesoporous metal oxides

can be further reduced to metal systems;

h) a "hard" template – that is a template which is itself a solid mesoporous or nanoporous

material. The voids of a preformed nano- or mesoporous solid, such as mesoporous silica

and nanoporous alumina, are impregnated with metal-containing precursor or precursors.

Subsequent solidification of these precursors and removal of the 'hard' template leads to

nano- or meso- porous catalysts. Such 'hard' templates may be produced using the

previously mentioned approaches (a-c) or may be purchased from a suitable supplier.

[0058] The In certain embodiments, the solidification process converts the metal precursor to a

chemical species which is a solid and which can self-support itself in the absence of the

templating agent. After this solidification process, the templating agent is removed by, for

example, washing the template out of the system.

[0059] The solidification process may involve a chemical transformation of a precursor by for

instance the oxidation or reduction of the metal precursor, or a change in the coordinating species

around the metal precursor with no change in formal oxidation state.

[0060] The reduction process can be carried out by any process known in the art by for example

(a) Electrochemically reducing – by poising the electrochemical potential of an electrode within

a suitable range so that the metal precursor is reduced to its corresponding metals on the surface

of the electrode; or (b) Chemically reducing - by adding a chemical reductant which reduces the

metal precursor to its corresponding metal. Suitable chemical reductants are for instance zinc,

iron, calcium or magnesium. Alternatively liquid reductants such as formaldehyde solutions;

formic acid solutions; or borohydride solutions; or gaseous reductants such as hydrogen,

formaldehyde, or boranes can be used.

[0061] Oxidation In certain embodiments, the oxidation process can be carried out by any

process known in the art for example (a) Electrochemically oxidising - by poising the

electrochemical potential of an electrode within a suitable range so that the metal precursor is

oxidised and forms an insoluble compound; or (b) Chemically oxidising – by adding a chemical

oxidant which oxidises the metal precursor to an insoluble compound. Suitable chemical

oxidants include solutions containing hydrogen peroxide or peroxide salts; perborates,

permanganates, chromates or chlorates; or gaseous oxidants such as ozone, chlorine or oxygen.

[0062] Another approach to cause solidification of the metal precursor involves the formation of

an insoluble chemical species. In one example, this can involve the addition of a compound

which changes the pH so that a metal oxide or hydroxide is precipitated. Compounds which can

be used to change the pH are acidic or basic materials, especially strong acids and bases for

instance sulphuric, nitric or hydrochloric acid or alkali metal hydroxides.

[0063] The metal precursor for the purposes of this invention should exhibit a high solubility in

the solution containing the templating agent. Such precursors can be the chloro complexes (i.e.

metal chlorides) or other halide complexes, or the amino (NH<sub>3</sub>), aqua (H<sub>2</sub>O), sulfito (SO<sub>3</sub>), or

cyano (CN) coordinated complexes, or complexes which contain a mixture of one or more of the

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above ligands. These complexes are particularly preferred for use in aqueous systems.

Complexes soluble in non-aqueous solvents include complexes containing acetylacetonate or

hexafluoroacetylacetonate, 1,10-phenanthroline and derivatives thereof, 1,5-cyclooctadiene and

derivatives thereof, or ethylenediamine and derivatives thereof.

[0064] The fifth aspect of the invention provides In a fifth aspect of the invention, there is

provided a process for the oxidation or reduction of organic and/or inorganic molecules

comprising contacting said molecules with the catalytic system of the first aspect or with the

catalyst of the third aspect.

[0065] The organic molecule can preferably comprise from one to about twelve carbon atoms,

preferably one to six carbon atoms, more preferably one to four carbon atoms. For example, the

organic molecule can be selected from methanol, formic acid, formate, methyl formate,

(CH<sub>3</sub>O)<sub>2</sub>CHO, or (CH<sub>3</sub>O)<sub>3</sub>CH or C1 molecules such as formaldehyde, and paraformaldehyde,

C2-aliphatic alcohols such as ethanol, 1,2-ethan-diol or partially oxidised C2 compounds e.g.

ethylene glycol; higher alkyl alcohols and related compounds or their oxidation products e.g.

isopropyl alcohol, propylene glycol etc. The organic molecule can also be a sugar, for instance

glucose, fructose, or sucrose.

[0066] The In certain embodiments, the oxidation of the organic and/or inorganic molecules by

the process of the fifth aspect allows the use of the catalytic system in a fuel cell wherein the

reactant is preferably a fuel for the fuel cell.

[0067] Preferred fuels for oxidation can be organic molecules, or mixtures of organic molecules.

Alternatively, the fuels can be inorganic molecules, for example hydrogen or hydrazine.

Mixtures of organic molecules and inorganic molecules can also be useful.

[0068] The In certain embodiments, the catalytic system of the first aspect may alternatively be

provided as a measuring element in a sensor. The sensor can be used to detect the organic and/or

inorganic molecules disclosed above.

[0069] In addition suitable analytes can be compounds which would not normally be fully

oxidised and which would not normally be suitable as fuels. Such suitable analytes can include

urea, thiourea, amino acids, peptides, proteins or carbohydrates.

[0070] The In certain embodiments, the catalyst or catalytic system described herein can also be

used to reduce an oxidant for example oxygen or hydrogen peroxide. The oxidant can be used

for example as a reactant for a fuel cell or as an analyte which is to be measured.

[0071] The sixth aspect of the invention provides In a sixth aspect of the invention, there is

provided an electrode comprising the catalytic system of the first aspect and/or the catalyst of the

third aspect. Said electrodes can be for use in an electrochemical cell, for example for use in a

fuel cell or a sensor.

[0072] The seventh aspect of the invention provides In a seventh aspect of the invention, there is

provided a fuel cell or sensor which comprises a catalytic system of the first aspect, a catalyst of

the third aspect or an electrode of the sixth aspect. Preferably, the fuel cell or sensor comprises

an electrode of the sixth aspect.

[0073] In particular, one or both of a pair of electrodes in a fuel cell or a sensor can comprise the

catalytic system of the first aspect and/or the catalyst of the third aspect. It will be appreciated

that further electrodes can also be present. Both of the electrodes must be fashioned on the

electrolyte such that they are in ionic contact with each other but not in electronic contact. For

example, if the electrolyte is a sheet (as is common in fuel cells), then the electrodes would be

placed on opposite sides of the sheet. In this example the configuration is used to minimise the

distance that any ion has to travel from one electrode to the other.

[0074] The configuration of a sensor can be similar to that of a fuel cell as described above. The

sensor can be used in detecting the presence or concentration of an organic or inorganic

molecule. The structure of a sensor can comprise for example two main electrodes in ionic

contact with each other through an electrolyte at least one of which comprises the catalytic

system of the invention. Extra electrodes may also be present to function as 'reference

electrodes' i.e. to provide a stable reference potential.

[0075] The eighth aspect of the invention provides In an eighth aspect of the invention, there is

provided a method for the detection of organic and/or inorganic molecules in a sample

comprising contacting said sample with a sensor as defined for the seventh aspect and detecting

the current due to oxidation or reduction of the molecules.

[0076] All preferred features of each of the aspects of the invention apply to all other aspects

mutatis mutandis.

[0077] The invention may be put into practice in various ways and a number of specific

embodiments will be described by way of example to illustrate the invention with reference to

the accompanying drawings, in which: drawings.

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Figure 1 shows SEM image and TEM image (inset) of Palladium mesoporous film electrochemically deposited on a gold foil at 0.16 V vs Ag/AgCl for 4 hr with a charge of 20 C cm<sup>-2</sup>.

Figure 2 shows high resolution scanning electron micrograph images of PtPdRuOs alloy chemically prepared from a liquid crystal phase, (A). Transmission electron micrograph (TEM) images of PtPdRuOs nanoparticles, (B). TEM images of a single PtPdRuOs particle, (C). Electron diffraction patterns of aggregated PtPdRuOs nanoparticles (D)

Figure 3 shows cyclic voltammograms for a mesoporous (a) and polycrystalline (b) Palladium disc microelectrode in 0.1 mol dm<sup>-3</sup>-HClO<sub>4</sub>+1 mol dm<sup>-3</sup>-methanol (dash line) and on a Nafion membrane equilibrated with 1 mol dm<sup>-3</sup>-methanol in water (solid line) at 20 °C (dE/dt: 50 mV s<sup>-1</sup>):

Figure 4 shows steady-state voltammograms for microelectrodes composed of mesoporous palladium, mesoporous PdPtRu and mesoporous PdPtRuOs catalysts on a Nafion<sup>TM</sup> membrane equilibrated with 1 mol dm<sup>-3</sup> CH<sub>3</sub>OH in water (a); and for mesoporous palladium on a Nafion<sup>TM</sup> membrane equilibrated with 0.15 mol dm<sup>-3</sup> HCOOH in water and 0.20 mol dm<sup>-3</sup> HCHO in water, (b).

Figure 5 shows steady-state voltammograms measured at 5 mV-s<sup>-1</sup> for a microelectrode containing mesoporous PtPdRuOs catalyst on a Nafion membrane equilibrated with 1 mol dm<sup>-3</sup> methanol in water at various temperatures. Inset: Potential dependence of mass activity of the quaternary alloy at 20 and 60 °C. Data taken from the positive-going scans.

Figure 6 shows a plot of limiting current versus methanol concentration (in water) for a mesoporous palladium catalyst/Nafion<sup>TM</sup> membrane catalyst system.

#### **EXAMPLES**

[0078] Example methods of preparing nanoporous or mesoporous Palladium-containing catalysts are described as follows:

[0079] Example 1: Electrochemical reduction method utilising a non-ionic surfactant that forms a liquid crystalline phase to prepare mesoporous palladium

[0080] The electroplating mixture consists of 12 wt % (NH<sub>4</sub>)<sub>2</sub>PdCl<sub>4</sub>, 47 wt % octaethylene glycol monohexadecyl ether (C<sub>16</sub>EO<sub>8</sub>), 2 wt % heptane and 39 wt % deionized water. The mixture was subjected to three heating-cooling cycles (temperature limits 20 and 80 °C) with vigorous shaking using a vortex mixer until a homogeneous mixture was formed. The mixture was then

allowed to stand at room temperature overnight. A conventional three-electrode system utilizing

a large-area platinum counterelectrode together with a saturated silver/silver chloride (Ag/AgCl)

was used to perform the electrodeposition. Mesoporous palladium films were produced by

electrochemical deposition from the electroplating mixture at +0.16 V vs Ag/AgCl onto a gold

film. The thickness of the palladium films were controlled by recording the total charge passed

during deposition. Typical deposition charge is about 20 C cm<sup>-2</sup>. Following deposition, the

palladium films were rinsed repeatedly in acetone and deionized water to remove impurities

(surfactant, electrolyte and organic solvents).

[0081] Low resolution SEM images of the film deposited on a gold foil show a smooth, dense

and uniform surface. High resolution SEM image exhibit uniform morphology characteristic of

leaf-like laminae as illustrated in figure 1A. The inset to Fig. 1(A) shows a TEM image of the

mesoporous film. The expected nanostructure is clearly seen with a pore diameter of around 3

nm and a pore center-pore center separation of around 6 nm. The specific electrochemical

surface area of the film measured using the underpotential Cu stripping technique (Chierchie, T.,

Mayer, C., Electrochim. Acta 33, 341 (1998)) is about 40 m<sup>2</sup> g<sup>-1</sup>. This value corresponding to a

surface area/volume ratio of 4.82x10<sup>8</sup> m<sup>2</sup> m<sup>-3</sup> is consistent with the hexagonal nanostructure of

the templated film.

[0082] Example 2: Electrochemical reduction method utilising a non-ionic surfactant that

forms a liquid crystalline phase to prepare Mesoporous Pd63Pt37 alloy:

[0083] Mesoporous Pd<sub>63</sub>Pt<sub>37</sub> alloy material was electroplated from a mixture consisting of 12 wt

% Na<sub>2</sub>PdCl<sub>6</sub>, 14 wt % H<sub>2</sub>PtCl<sub>6</sub>, 26 wt % H<sub>2</sub>O, 2 wt % heptane, 47 wt % surfactant C<sub>16</sub>EO<sub>8</sub>.. The

mixture was subjected to three heating-cooling cycles (temperature limits 20 and 80 °C) with

vigorous shaking using a vortex mixer until a homogeneous mixture was formed. The mixture

was then allowed to stand at room temperature overnight. A conventional three-electrode system

utilizing a large-area platinum counterelectrode together with a saturated silver/silver chloride

(Ag/AgCl) was used to perform the electrodeposition. The Palladium alloy mesoporous film was

deposited onto a gold film at a potential of 0.06 V vs Ag/AgCl. Following deposition, the film

was rinsed repeatedly in acetone and deionized water to remove impurities (surfactant,

electrolyte and organic solvents).

[0084] Example 3: Chemical reduction method utilising a non-ionic surfactant that forms a

liquid crystalline phase to produce mesoporous PtPdRuOs alloy:

[0085] A mixture containing 0.364 g Na<sub>2</sub>PdCl<sub>6</sub> (Aldrich), 0.50 g H<sub>2</sub>PtCl<sub>6</sub>.xH2O (Aldrich), 0.202 g RuCl<sub>3</sub>.xH2O (Aldrich), 0.30 g Na<sub>2</sub>OsCl<sub>6</sub>.xH2O (Aldrich), 1.265g Water, 1.88 g octaethylene glycol monohexadecyl ether non-ionic surfactant (C16EO8,Fluka) was homogeneously mixed with vigorous shaking using a vortex mixer and subjected to three heating-cooling cycles (temperature limits 20 and 80 °C) before being allowed to stand at room temperature for 12 hours. The quaternary catalyst was then precipitated by adding pieces of metallic Zn(Aldrich) to the reactant mixture to chemically reduce the metal salts present in the aqueous domains of the mixture. This reduction process was allowed to continue at room temperature until the mixture became fully black. The black deposit was washed successively with acetone, hydrochloric acid, water and acetone. The recovered product was then dried in an oven at 80 °C.

[0086] Figure 2 shows high resolution scanning electron micrograph images of the PtPdRuOs alloy (A). Transmission electron micrograph (TEM) images of PtPdRuOs nanoparticles, (B) and TEM images of a single PtPdRuOs particle showing the well developed porous structure, (C). Electron diffraction patterns of aggregated PtPdRuOs nanoparticles (D)

# [0087] Example 4: Chemical reduction method utilising a hard template to produce mesoporous palladium

[0088] The template synthesis of mesoporous palladium utilising a 'hard' template can be carried out as follows: 5 g aqueous solution of 1 mol dm<sup>-3</sup> Pd(NH<sub>3</sub>)<sub>4</sub>Cl<sub>2</sub> (Aldrich) was impregnated into 1 g calcined mesoporous silica SBA-12 powder. After being completely dried at 100 °C, the impregnation was repeated three times. Then the SBA-12 sample impregnated with Pd(NH<sub>3</sub>)<sub>4</sub>Cl<sub>2</sub> was transferred into a glassy tube reactor with two fritted disks. The palladium compound was reduced to Pd using H<sub>2</sub> flowing through the tube while the temperature was increased from 20 °C to 300 °C at 1 °C/min, and then maintained at 300 °C for 2 h. The resultant sample was exposed to air after the tube reactor was cooled to room temperature. The Pd/SBA-12 sample was digested in 3 mol dm<sup>-3</sup> HF solution for 2 h in order to dissolve our the SBA-12. The black product was centrifugatedly filtered, washed with distilled water and dried in a vacuum oven at about 60 °C.

# [0089] Example 5: Chemical reduction method utilising a hard template to produce mesoporous palladium

[0090] In this approach silica powder is used as the hard template and Palladium is supplied as an organo-metallic compound in a fluidized bed reactor using a metal-organic chemical vapor deposition (MOCVD) technique (Jean-Cyrille Hierso, Roselyne Feurer, and Philippe Kalck, Chem. Mater. 2000, 12, 390-399). Four grams of silica powder (sieved between 100 and 200 μm) is supported on a stainless steel mesh with 25 μm openings above a sintered glass frit of supporting the complex (0.4)of (porosity 5) precursor  $(\eta_3$ -Allyl)(hexafluoroacetylacetonato)palladium). The reactor was evacuated to 0.1 Torr and kept at this pressure for 1 h while the silica powder was heated to 100 °C by ethylene glycol passing through a double envelope in the reactor wall. After this time, the silica powder was heated to 110 °C (the deposition temperature) and the pressure inside the reactor was raised to 30 Torr by the addition of helium (helium flow: 86 mL min<sup>-1</sup>) at the bottom of the reactor. This pressure was determined so that the velocity of gas through the fluidized bed was approximately four times the minimum bubbling velocity of the powder. The lower portion of the reactor was then heated to 80 °C to sublime the complex into the stream of helium, and a stream of hydrogen was introduced just before the fluidized bed. After 2.5 h, the heating was stopped and the reactor allowed to cool to room temperature. The pressure was returned to atmospheric pressure, and the silica powder was recovered. The Pd/silica sample was washed with 3 mol dm<sup>-3</sup> HF solution for 2 h. The black product was filtered under centrifugation, washed with distilled water and dried in a vacuum oven at about 60 °C.

# [0091] Example 6: Chemical reduction method utilising the sol-gel route to produce mesoporous palladium

[0092] 1 g of the P-123 (BASF) triblock copolymer was dissolved in 10 g butanol with stirring for 30 min. To this solution, 0.005 mol dm<sup>-3</sup> PdCl<sub>2</sub> was added with vigorous stirring for 30 min. The resulting sol solution was gelled in an open dish at 60 °C in air for one week. The aged gel sample was then calcined in air at 500 °C for 4 h to remove the block copolymer surfactant species. The resultant palladium oxide was transferred into a glass tube reactor with two fritted disks after it cooled down. The palladium compound was reduced to Pd with H<sub>2</sub> flowing through the tube while the temperature was increased from 20 °C to 300 °C at 1 C/min, and then maintained at 300°C for 2 h. The resultant sample was exposed to air after the tube reactor was cooled to room temperature.

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# [0093] Example 7: Chemical reduction method utilising the microemulsion route to produce mesoporous palladium

[0094] Microemulsions containing the metallic salts were prepared by adding 1 mol dm<sup>-3</sup> H<sub>2</sub>PdCl<sub>6</sub> to 0.2 mol dm<sup>-3</sup> Brij<sup>®</sup>76 in hexane to obtain a ratio of 2/1 H<sub>2</sub>O/surfactant in the emulsion. Under these conditions the final concentration of the palladium salt is around 5×10<sup>-3</sup> mol dm<sup>-3</sup> in the microemulsion In the same way, a microemulsion containing the reductant was prepared by adding an aqueous solution of hydrazine hydroxide to 0.2 mol dm<sup>-3</sup> Brij<sup>®</sup>76 solution in hexane. The amount of hydrazine added was adjusted to produce a concentration of 0.05 mol dm<sup>-3</sup> in the microemulsion. Mixing the two microemulsions produces a colloidal suspension of palladium nanoclusters. The black suspension was filtered under centrifugation, and then washed subsequently with ethanol and water. Finally, the solid product was dried at 80°C.

[0095] Example methods of preparing electrode assemblies are described as follows:

### [0096] Example 8: Direct mechanical assembly

[0097] A conductive electrode with an electrodeposited mesoporous palladium or palladium containing film (for instance prepared by process 1 or 2 above) was pressed into direct contact with a Nafion<sup>TM</sup> membrane. Utilisation of a small amount of soluble Nafion<sup>TM</sup> solution on the electrode surface before the interface is formed followed by curing at 120°C aids the long-term stability of the interface.

### [0098] Example 9: Ink route preparation of mesoporous electrode-membrane assembly

[0099] A thin catalyst layer film was sprayed from the inks consisting of mesoporous Pd, 5 wt.% solubilized Nafion®(supplied by DuPont Company), dionized (DI) water, (NH<sub>4</sub>)<sub>2</sub>C<sub>2</sub>O<sub>4</sub> (pore forming additive) and isopropanol. The ratio of mesoporous palladium catalyst to Nafion® is 3:1. The resultant ink was then sprayed by air-brushing over XC-72 carbon coated carbon papers to produce the electrode. After the ink on the carbon paper was dried up, extra Nafion<sup>TM</sup> solution in iso-propanol was sprayed on the surface of the electrode. The membrane electrode assembly was prepared by hot pressing the electrode onto a Nafion<sup>TM</sup> 117 membrane at 140 °C and 200 atm for 1.5 min.

## [0100] Example 10: Decal approach for producing catalyst layer

[0101] An ink as described (4) was prepared and uniformly painted on a ptfe decal blank. After the applied inks were dried, the thin film was transferred from the ptfe decal blank to the Na<sup>+</sup> form of Nafion<sup>®</sup> 115 membrane by hot-pressing at 160–200 °C and 5–9 MPa for 150 s. After the Page 19 of 27

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decal blanks were removed, the hydrophilic film/Nafion<sup>®</sup> membrane assemblies were ion-exchanged to the protonated form by immersing them in lightly boiling 0.5 M H<sub>2</sub>SO<sub>4</sub> solution for about 1 h.

# [0102] Example 11: Nafion<sup>TM</sup> impregnation of film electrode

[0103] An ink was prepared containing ethylene glycol (10wt%), ptfe suspension (10wt% ptfe, T30, DuPont), and mesoporous palladium. After mechanical mixing and agitation in an ultrasound bath, the mixture was brushed over a carbon paper/XC72R carbon diffusion layer. The mixture was dried in an oven at 80°C and then sintered at 360°C for 20 minutes.. After allowing the electrode to cool, it was brushed with 5 wt% Nafion<sup>TM</sup> in alcohol solution (Aldrich) and this solution was allowed to diffuse into the inner pores of the electrode film for ten minutes at room temperature, followed by drying in air at 80 °C to eliminate residual aliphatic alcohols. Then the electrode was then hot-pressed onto Nafion<sup>TM</sup> 117 membrane at 160–200 °C and 5–9 MPa for 150 s..

[0104] Example results of performance of electrode assemblies

# [0105] Example 12: Superior performance of mesoporous or nanoporous palladium containing catalyst in contact with an ion-exchange electrolyte

[0106] This example shows that when a mesoporous or nanoporous catalyst is in contact with an ion exchange membrane the activity of the system is greater than seen if either of these separate aspects (meso/nano porosity and ion exchange electrolyte) are absent. The mesoporous palladium electrode was produced using as set out in Example 1 and it was assembled according to Example 8. In figure 3 methanol oxidation at both mesoporous and polycrystalline palladium electrodes in both an aqueous electrolyte (0.1 mol dm<sup>-3</sup> HClO<sub>4</sub>) and at a exchange membrane (Nafion<sup>TM</sup>) are displayed. The currents are normalised to the total surface area of the catalyst. When the aqueous electrolyte (0.1 mol dm<sup>-3</sup> HClO<sub>4</sub>) is used (dotted lines) a similar response during methanol oxidation is seen independent of the catalyst morphology – namely a sharp anodic peak during the positive-going scan that quickly decays to a low-level current at higher potentials and a similar peak, although shifted in potential, during the negative-going scan. The height of the forward and reverse anodic peaks are very close. However, when Nafion<sup>TM</sup> is used as the electrolyte, the response is dependent upon the morphology of palladium used. At polycrystalline palladium electrodes, methanol oxidation commences at about 0.75 V, and a sharp anodic peak followed by current decay are seen during the positive-going scan i.e. similar

to the response seen for polycrystalline palladium in HClO<sub>4</sub>. During the reverse scan only a small anodic peak is produced. In contrast, at the mesoporous palladium electrode, the onset potential of the methanol oxidation is shifted negative by about 150 mV. This represents a significant improvement in performance. Furthermore, the rate of methanol oxidation (i.e. the magnitude of the oxidation current) when the mesoporous catalyst is used is about five times larger than that on the polycrystalline palladium. The rate of methanol oxidation is so high on the mesoporous palladium that a current plateau is seen over 0.85~1.30 V. Such a plateau is also evident in slow-scan voltammograms at 2 mV s<sup>-1</sup> and strongly suggests that the electrode reaction is diffusion controlled at high overpotential. Thus a significant enhancement in catalytic activity is seen when mesoporous palladium is used in contact with a solid polymer electrolyte.

# [0107] Example 13: Oxidation of a range of organic species using catalyst comprising mesoporous or nanoporous palladium in contact with an ion-exchange electrolyte

<u>[0108]</u> This example shows the performance of mesoporous/nanoporous palladium comprising catalyst in contact with an ion exchange membrane towards the oxidation of various organic compounds and the beneficial effect on activity obtained upon addition of other compounds to the nanoporous/mesoporous palladium catalyst. The results are presented in Figure 4. The nanoporous/mesoporous palladium catalyst was prepared according to Example 1. The nanoporous/mesoporous PdPtRu was produced according to Example 2 except that some RuCl<sub>3</sub> was added to the templating solution. The PdPtRuOs was produced according to the method described in Example 3. In all cases the electrode assemblies were produced using the 'direct mechanical assembly' method (above). The ion exchange electrode used was a Nafion<sup>TM</sup> 117 membrane.

[0109] In figure 4(a) the performance of the mesoporous palladium/Nafion system is examined for the oxidation of three different organic compounds of interest in fuel cells and sensors: methanol, formaldehyde and formic acid. only for the oxidation of formaldehyde and formic acid at this kind of composite electrode. The onset potential for the limiting current is around 0.80 and 0.30 V for the oxidation of formaldehyde and formic acid respectively. Therefore, the composite electrode is applicable to formaldehyde and formic acid sensors. The difference in onset potentials above suggests that the sensor electrode has high selectivity for the detection of formaldehyde and formic acid. In all cases a diffusion limiting current or current plateau is found indicating that the rate of oxidation becomes limited by the rate of reactant transport to the

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catalyst. The low potential at which formic acid oxidation occurs (the reaction commences at around 0 V), and the exceptionally low potential at which the diffusion limited current occurs (0.3V) indicate that formic acid can be decomposed to CO<sub>2</sub> at potentials relevant to fuel cell operations. Mesoporous Palladium is therefore a highly efficient anode electrocatalyst for an ion exchange electrolyte-based formic acidic fuel cell. The mesoporous palladium/Nafion catalytic system is also active for the oxidation of formaldehyde and methanol as shown in the other traces in Figure 4(a). For both of these other fuels, a limiting current is also observed. These limiting currents are at different values from that seen from formic acid due to the different transport properties and concentration of each of the species being studied.

[0110] In Figure 4(b), the beneficial effect of combining the nanoporous/mesoporous palladium catalyst with other compounds on the oxidation of methanol is described. In all cases a diffusion limiting current is obtained at a suitably high potential, as evidenced by the plateau in activity. This indicates that the performance is limited by the rate at which methanol can reach the catalyst surface and not by the rate of reaction. When platinum and ruthenium is alloyed with the palladium, the activity of the catalyst increases significantly, as shown by the -220mV shift in the onset potential between the mesoporous palladium and the mesoporous PdPtRu. When Osmium is further added as an alloying element, the onset potential shifts a further -150mV, indicating a further significant improvement in activity. The quaternary PtRuPdOs/ion exchange electrolyte catalytic system appears to be more active than commercially obtained PtRu catalyst for the methanol oxidation reaction.

### [0111] Example 14: Catalytic system for inclusion within a direct methanol fuel cell

**[0112]** Further evidence of the efficacy of the PtRuPdOs/ion exchange electrolyte catalytic system for the oxidation of methanol is provided in Figure 5 which shows the performance of the mesoporous Pd<sub>27</sub>Pt<sub>38</sub>Ru<sub>23</sub>Os<sub>12</sub>/Nafion catalytic system as a function of temperature. Increasing the temperature from 20°C to 60°C leads to a shift of -100mV in the onset potential. Increasing the temperature also increases the limiting current, as higher temperatures improve the mass transport characteristics of methanol in the ion exchange electrolyte. Displayed inset of Figure 5 is the mass activity of the catalyst. At 0.3 V, and activity of over 30 A g<sup>-1</sup> is achievable at 60°C.

### [0113] Example 15: Preparation of a methanol sensor

[0114] Owing to a linear relation between the diffusion-limiting current and the concentration of active reactants, the nanoporous/mesoporous palladium catalytic system can be directly used as a

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methanol sensor. This type of sensor avoids the need for an aqueous electrolyte. The sensor incorporates the catalytic system and a method to pass the analyte containing the unknown concentration of methanol over the catalytic system. The response of the limiting current to variations in the concentration of methanol is determined by the sorption isotherm of methanol from the feed solution into the electrolyte. Methanol is more likely to dissolve in the microvoids of the membrane rather than its polytetrafluoroethylene backbone. If the sorption of methanol into the microvoids dominates, a Langmuir sorption isotherm would be expected.

[0115] The dependence of the limiting current upon the methanol concentration of the feed solution can be given as:

$$I_d = \frac{I_d^{\text{max}} kc}{1 + kc} \tag{1}$$

where  $I_d$  and  $I_d^{max}$  are the diffusion limited current at a methanol concentration c and at the maximum concentration of methanol in the membrane. k is the Langmuir affinity constant. Thus it is possible to calculate an unknown methanol concentration in water by measuring the limiting current and solving equation (1) for c, once the parameters  $I_d^{max}$  and k are known for a specific configuration.

[0116] Displayed in figure 6 is a plot of limiting current versus methanol concentration (in water) for a mesoporous palladium catalyst/Nafion<sup>TM</sup> membrane catalyst system. This catalyst system is the same as that used in Example 13. The limiting current increases with methanol concentration in water reaching a maximum at about 2 mol dm<sup>-3</sup> of methanol in water. Fitting the curve in Fig. 6 gives  $I_d^{max}$ =3.36  $\mu$ A and k=2.0 with a confidence of 98.5 %. Thus this system could now be used to measure the unknown methanol concentration in water by measuring the limiting current and solving equation (1) for c using the values of  $I_d^{max}$  and k displayed above.

**CLAIMS** 

(Original) A catalytic system comprising a catalyst comprising nanoporous or 1.

mesoporous palladium and an ion-exchange electrolyte.

2. A catalytic system as claimed in claim 1 wherein the catalyst further

comprises one or more of platinum, gold, ruthenium, rhodium, osmium, iridium, silver, nickel,

copper, cobalt, iron, chromium, lead, vanadium or tungsten.

3. (Original) A catalytic system as claimed in any one of claims 1 to 2 wherein the catalyst

is Pd, PdPt, PdAu, PdPtRu, PdPtIr, PdPtAu, PdPtRuIr, PdPtRuOs or PdPtRuIrOs.

4. (Currently Amended) A catalytic system as claimed in any one of claims 1 to 3 to 2

wherein the ion-exchange electrolyte is a cation exchange electrolyte.

5. (Original) A multi-component catalyst comprising a nanoporous or mesoporous

palladium and one or more of platinum, gold, ruthenium, rhodium, osmium, iridium, silver,

nickel, copper, cobalt, iron, chromium, lead, vanadium, tungsten, carbon, nitrogen, oxygen,

sulphur, selenium, tellurium or phosphorous.

6. (Original) A process for the production of a catalyst as claimed in claim 5, said process

comprising solidifying one or more metal precursors in the presence of a templating agent and

then removing the templating agent.

7. (Currently Amended) A process for the oxidation or reduction of inorganic and/or

organic molecules comprising contacting said molecules with the catalytic system as claimed in

any one of claims claim 1 to 4 or the catalyst as claimed in claim 5 or 2, or a multi-component

catalyst comprising a nanoporous or mesoporous palladium and one or more of platinum, gold,

ruthenium, rhodium, osmium, iridium, silver, nickel, copper, cobalt, iron, chromium, lead,

vanadium, tungsten, carbon, nitrogen, oxygen, sulphur, selenium, tellurium or phosphorous.

(Original) A process as claimed in claim 7 wherein the organic molecule has from one 8.

to twelve carbon atoms.

9. (Currently Amended) A process as claimed in claim 7 or 8 wherein the organic

molecule is methanol.

10. (Cancelled)

11. (Currently Amended) A process for the manufacture of a catalytic system as claimed in

any one of claims 1 to 4 claim 1 or 2 comprising contacting the catalyst with the ion-exchange

electrolyte.

12. (Currently Amended) An electrode comprising a catalytic system as claimed in any one

of claims claim 1 to 4 or a catalyst as claimed in claim 5 or 2, or a multi-component catalyst

comprising a nanoporous or mesoporous palladium and one or more of platinum, gold,

ruthenium, rhodium, osmium, iridium, silver, nickel, copper, cobalt, iron, chromium, lead,

vanadium, tungsten, carbon, nitrogen, oxygen, sulphur, selenium, tellurium or phosphorous.

13. (Original) A fuel cell comprising an electrode as claimed in claim 12.

14. (Original) A sensor comprising an electrode as claimed in claim 12.

15. (Original) A method for the detection of organic and/or inorganic molecules in a sample

comprising contacting said sample with a sensor as claimed in claim 14 and detecting the current

due to the oxidation or reduction of the molecules.

Claims 16-21 (Cancelled)

22. (New) The catalytic system of claim 3 wherein the ion-exchange electrolyte is a cation

exchange electrolyte.

23. (New) The process of claim 7 wherein, in the catalytic system, the catalyst is Pd, PdPt,

PdAu, PdPtRu, PdPtIr, PdPtAu, PdPtRuIr, PdPtRuOs or PdPtRuIrOs.

24. (New) The process of claim 7 wherein, in the catalytic system, the ion-exchange

electrolyte is a cation exchange electrolyte.

25. (New) The process of claim 8 wherein the organic molecule is methanol.

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26. (New) The process of claim 11 wherein, in the catalytic system, the catalyst is Pd, PdPt,

PdAu, PdPtRu, PdPtIr, PdPtAu, PdPtRuIr, PdPtRuOs or PdPtRuIrOs.

(New) The process of claim 11 wherein, in the catalytic system, the ion-exchange 27.

electrolyte is a cation exchange electrolyte.

28. (New) The electrode of claim 12 wherein, in the catalytic system, the catalyst is Pd, PdPt,

PdAu, PdPtRu, PdPtIr, PdPtAu, PdPtRuIr, PdPtRuOs or PdPtRuIrOs.

29. (New) The electrode of claim 12 wherein, in the catalytic system, the ion-exchange

electrolyte is a cation exchange electrolyte.

30. (New) The fuel cell of claim 13 wherein, in the electrode comprising the catalytic system,

the catalyst is Pd, PdPt, PdAu, PdPtRu, PdPtIr, PdPtAu, PdPtRuIr, PdPtRuIr, PdPtRuIrOs.

(New) The fuel cell of claim 13 wherein, in the electrode comprising the catalytic system, 31.

the ion-exchange electrolyte is a cation exchange electrolyte.

32. (New) The sensor of claim 14 wherein, in the electrode comprising the catalytic system,

the catalyst is Pd, PdPt, PdAu, PdPtRu, PdPtIr, PdPtAu, PdPtRuIr, PdPtRuOs or PdPtRuIrOs.

33. (New) The sensor of claim 14 wherein, in the electrode comprising the catalytic system,

the ion-exchange electrolyte is a cation exchange electrolyte.

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**ABSTRACT** 

Nanoporous/mesoporous palladium catalyst

The present invention provides a catalytic system comprising a catalyst comprising nanoporous

or mesoporous palladium and an ion-exchange electrolyte, processes for manufacturing the

catalytic system and catalyst, and processes for oxidising or reducing organic and/or inorganic

molecules using the catalyst or catalytic system.

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